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ESCMID COVID-19 Living guidelines: drug treatment and clinical management

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Journal Pre-proof

ESCMID COVID-19 Living Guidelines: Drug Treatment and Clinical Management

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Abstract

Scope: In January 2021, the ESCMID Executive Committee (EC) decided to launch a new initiative to develop ESCMID guidelines on several COVID19-related issues, including treatment of COVID-19.

Methods: An ESCMID COVID-19 guidelines task force was established by the ESCMID Executive Committee. A small group was established, half appointed by the chair, and the remaining selected with an open call. Each panel met virtually once a week. For all decisions, a simple majority vote was used. A long list of clinical questions using the PICO (population, intervention, comparison, outcome) format was developed at the beginning of the process. For each PICO, two panel members performed a literature search with a third panelist involved in case of inconsistent results. Voting was based on the GRADE approach.

Questions addressed by the guideline and recommendations: A synthesis of the available evidence and recommendations are provided for each of the 15 PICOs, which cover use of hydroxychloroquine, bamlanivimab alone or in combination with etesevimab, casirivimab combined with imdevimab, ivermectin, azithromycin and empirical antibiotics, colchicine, corticosteroids, convalescent plasma, favipiravir, remdesivir, tocilizumab, and interferon β-1a, as well as the utility of antifungal prophylaxis and enoxaparin. In general, the panel recommended against the use of hydroxychloroquine, ivermectin, azithromycin, colchicine, and interferon β-1a. Conditional recommendations were given for the use of monoclonal antibodies in highrisk outpatients with mild-moderate COVID-19, and remdesivir. There was insufficient evidence to make a recommendation for use of favipiravir and antifungal prophylaxis, and it was recommended that antibiotics should not be routinely prescribed in patients with COVID-19 unless bacterial coinfection or secondary infection is suspected or confirmed. Tocilizumab and corticosteroids was recommended for treatment of severe COVID-19 but not in outpatients with non-severe COVID-19.

Scope

The aim of the present guidance is to provide evidence-based recommendations for management of adults with coronavirus disease 2019 (COVID-19). More specifically, the goal is to aid clinicians managing patients with COVID-19 at various levels of severity including outpatients, hospitalized patients, and those admitted to intensive care unit (ICU). Considering the composition of the panel, mostly clinical microbiologists or infectious disease specialists with no pulmonology or intensive care background, we focus only on pharmacological treatment and do not give recommendations on oxygen supplement/support. Similarly, as no pediatricians were included in the panel, the recommendations are only for adult patients with COVID-19. Considering the current literature, no guidance was given for special populations such as the immunocompromised.

Background

The pandemic caused by severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) has had a dramatic impact on healthcare systems, the global economy, and social life. The clinical spectrum of COVID-19 induced by SARS-CoV-2 is broad with the majority of infected individuals experiencing only mild or subclinical illness, especially in the early phase of disease [1]. However, 14-30% of hospitalized patients with COVID-19 develop severe respiratory failure requiring intensive care [2-4]. Additionally, as the angiotensine-converterend enzyme 2 (ACE2) receptor is widely distributed in human organs and tissues, manifestations of COVID-19 involve many organs including the central nervous system, kidneys, myocardium, and gut.

As of July 6, 2021, worldwide more than 184 million people have tested positive for SARS-CoV-2 and nearly 4 million have died of COVID-19. In light of this dramatic situation, the ongoing pandemic generated a historical effort involving many researchers worldwide and prompted an unprecedented number of clinical trials. According to ClinicalTrials.gov, as of March 10, 2021, nearly 5000 studies are investigating COVID-19.

Motivations for guideline development

ESCMID did not develop its own recommendations at the start of the pandemic for several reasons: clinical overload of most members, avoid duplication of ongoing

efforts, heterogeneity of national recommendations, and lack of appropriate evidence. The latter is particularly relevant, since issuing guidance based on inappropriate evidence-base might do more harm than good. In January 2021, the ESCMID Executive Committee (EC) decided to launch a new initiative to develop ESCMID guidelines on several COVID19-related issues.

Methods

An ESCMID COVID-19 guidelines task force was established by the ESCMID EC. For each set of guidelines, a small group was established (10-15 panelists). Half were appointed by the chair, in agreement with the EC, and the remaining were selected with an open call carried out on January 2021 and advertised on all ESCMID channels. The ESCMID guidelines subcommittee evaluated the applications and issued a recommendation about inclusion/exclusion of each applicant. As for all ESCMID initiatives, balance in terms of gender, clinical specialty, and country was maintained.

Project management

Each panel met virtually once a week. For all decisions, a simple majority vote was used and a decision was made in case of ≥80% of agreement.

. A long list of clinical questions using the PICO (population, intervention, comparison, outcome) format was developed at the beginning of the process. A maximum number of 15 PICOs was set and selected by vote (the 15 top-rated PICOs were chosen). Criteria for prioritization and vote were general interests by clinicians with clinical microbiology and infectious disease background and availability of evidence, especially for critical outcomes that included mortality or disease progression [ICU admission or need for mechanical ventilation or extracorporeal membrane oxygenation (ECMO)]. Additional PICOs will be developed at a further stage.

Evidence review

To avoid duplication of efforts, rather than performing a systematic review of the literature for each PICO, each panel reviewed whether evidence for each PICO was already available among the many ongoing initiatives [6-8]. For each PICO and evidence synthesis, ADOLOPMENT criteria were used (Table 1). For each PICO, two panel members performed a literature search with a third panelist involved in case of

inconsistent results. The results of the searches were presented to the panel during weekly meetings for discussion and voting (quality of evidence, evidence-to-decision criteria, need for update, etc.) based on the GRADE approach.

Definitions

WHO severity criteria for COVID-19 were used [9]. Data from the European Center for Disease Prevention and Control (ECDC) was used to define risk factors and groups for severe COVID-19 [10].

Questions addressed by guidelines and recommendations

For each PICO question, the motivations for use, patient preferences and additional comments are presented in Supplementary Appendix 1. A summary of all recommendations is presented in Table 2.

What is the effect of hydroxychloroquine treatment on mortality or disease progression in patients with mild COVID-19 compared with no treatment?

Narrative synthesis of evidence

Twenty-three randomized trials in >10,000 patients have assessed the effect of hydroxychloroguine (HCQ) on COVID-19 compared with standard of care (SOC). For the present assessment, 19 trials were included (Table 3). HCQ had no impact on death (risk ratio [RR] 1.06, 95% confidence interval [CI] 0.97-1.16) or need for mechanical ventilation (RR 1.08, 95% CI 0.91 to 1.28). The majority of patients were included in the RECOVERY and SOLIDARITY trials. RECOVERY is an investigatorinitiated platform trial at 176 hospitals in the UK. Within this, 1561 patients were randomized to receive HCQ and 3155 to SOC. No difference in 28-day mortality was observed between HCQ and SOC (RR 1.09; 95% CI, 0.97-1.23; p=0.15) [11]. In SOLIDARITY, hospitalized patients with COVID-19 were randomized to remdesivir (n=2750), HCQ (n=954), lopinavir (n=1411), interferon β -1a (n=2063), or SOC (n=4088). The primary outcome was 28-day mortality and occurred in 104 of 947 patients receiving HCQ and in 84 of 906 controls (RR 1.19; 95% CI 0.89–1.59; p=0.23) [12]. HCQ was not effective in smaller randomized controlled trials (RCTs) in hospitalized patients [13-15], hospitalized patients with severe [16-18] or mildmoderate disease [19-25], or outpatients [26-28]. Lastly, HCQ has not been

associated with a faster decline of viral load or higher virological cure compared to SOC in hospitalized patients [19, 22, 25, 28, 29].

Safety

Concerns for safety and potential harm have been raised in observational trials and RCTs evaluating patients receiving HCQ. RECOVERY reported that those receiving HCQ experienced longer in-hospital stay, lower probability of being discharged alive within the 28-day study period (RR 0.92; 95% CI 0.85–0.99), and higher chance to receive mechanical ventilation (30.7% vs. 26.9%; RR 1.14; 95% CI 1.03–1.27). A trend towards greater harm with HCQ was also seen in SOLIDARITY and other RCTs [12].

Recommendation

Strong recommendation against use of HCQ for COVID-19 [quality of evidence (QoE): high for critical outcomes].

What is the effect of bamlanivimab alone or in combination with etesevimab in reducing the risk of disease progression or mortality in patients with mild COVID-19 compared with no treatment?

Narrative synthesis of evidence

Bamlanivimab and etesevimab are recombinant neutralizing human IgG1k monoclonal antibodies (mAb) directed against the spike protein of SARS-CoV-2. They were evaluated in BLAZE-1, a randomized, double-blind, placebo-controlled, multipart phase 2/3 trial enrolling outpatients with COVID-19. In the first dataset of BLAZE-1, bamlanivimab showed a trend towards decreased viral load vs. placebo with a significant difference for the 2800 mg dose. [30]. The second dataset of the BLAZE-1 trial analyzed patients randomized to receive a single infusion of bamlanivimab at different dosages, combined bamlanivimab and etesevimab, or placebo. Compared with placebo, a significant decrease in viral load was observed only for combination treatment [log -0.57 (95% CI, -1.00—0.14; p=\(\Delta 0.01 \)]. The percentages of patients with COVID-19—related hospitalizations or emergency department visits was 5.8% (n=9) for placebo, 1.0% (n=1) for 700 mg, 1.9% (n=2) for 2800 mg, 2.0% (n=2) for 7000 mg, and 0.9% (n=1) for combination treatment [31].

On March 10, 2021, via press release, a new analysis on 769 high-risk patients with mild to moderate COVID-19 receiving bamlanivimab plus etesevimab (n=511) or placebo (n=258) was presented. There were four hospitalizations in patients taking bamlanivimab and etesevimab compared to 15 for placebo (risk reduction 87%; p<0.0001) [32].

Overall, in high-risk outpatients bamlanivimab alone (RR 0.26; 95% CI 0.09–0.75; Table 4) or combined with etesevimab (RR 0.30; 95% CI 0.16–0.59; Table 5) is associated with reduced hospitalization. Bamlanivimab plus etesevimab is also associated with reduction in 29-day mortality (RR 0.05; 95%CI 0.00–0.80) in the same population (Table 5).

Bamlanivimab was effective in preventing severe disease among residents and staff of long-term care facilities (BLAZE-2 trial) [33], but not in recovery of hospitalized patients [34].

In vitro studies suggest that bamlanivimab plus etesevimab retains in vitro susceptibility to the B.1.1.7 (Alpha – UK variant), but has markedly reduced activity against the P1 (Gamma, Brazilian) and B.1.351 (Beta, South African) variants. Lastly, the SARS-CoV-2 variant B.1.617 (Delta, Indian) seems to be resistant to bamlanivimab, but its activity may be restored when combined with etesevimab.

Safety

Infusion-related adverse events were reported in 14% of patients in one study. Overall, adverse events were not higher vs. placebo in all studies [30-32].

Recommendation

Weak recommendation against use of bamlanivimab alone (QoE: very low).

Conditional recommendation for use of bamlanivimab plus etesevimab in high-risk outpatients with mild to moderate COVID-19 (QoE: moderate).

What is the effect of casirivimab combined with imdevimab in reducing the risk of disease progression or mortality in patients with mild COVID-19 compared with no treatment?

Narrative synthesis of evidence

Casirivimab and imdevimab were assessed in a phase 1-3 trial in which patients were randomized to placebo, 2.4 g of combination therapy (casirivimab 1200 mg and imdevimab 1200 mg), or 8.0 g of combination therapy (4.0 g casirivimab and 4.0 g imdevimab). The combination of casirivimab and imdevimab was significantly associated with reduction of viral load [35], COVID-19–related hospitalization, and all-cause death vs. placebo (71.3% reduction; 1.3% vs. 4.6%; p<0.0001) [35]. A significant effect was also seen in patients with baseline positive serum anti-SARS-CoV-2 antibodies [35]. Casirivimab combined with imdevimab was associated with a lower rate of hospitalization (RR 0.27; 95% CI 0.11–0.65; Table 6).

Hospitalized patients

The combination of casirivimab (4.0 g) plus imdevimab (4.0 g) was assessed in RECOVERY and was associated with lower 28-day mortality among anti-SARS-CoV-2 Ab seronegative patients at baseline (RR 0.80; 95% CI 0.70–0.91; p=0.0010) [36].

Safety

The rate of adverse events was similar between patients receiving casirivimab plus imdevimab or placebo, while the combination showed fewer serious adverse events [35, 36].

Recommendation

Conditional recommendation for use of combination casirivimab plus imdevimab in high-risk outpatients with mild-moderate COVID-19 (QoE: moderate for hospitalization; low for 29-day mortality).

What is the effect of ivermectin in reducing the risk of disease progression or mortality in patients with mild COVID-19 compared with no treatment?

Narrative synthesis of evidence

Ivermectin has been evaluated in 18 RCTs using different dosing regimens and number of doses (1 to 5). Ten studies primarily had a virological outcome, i.e. virological reduction or clearance [37-46], while most reported secondary clinical outcomes like mechanical ventilation and death. Overall, 11 studies showed a positive effect of ivermectin while 7 did not (Table 7), with the largest reporting no effects [47,

48]. The committee was thus uncertain whether ivermectin increased or decreased the chance of need for mechanical ventilation or death.

Safety

While no serious adverse events were recorded (Table 7), there was uncertainty with regards to adverse events and gastrointestinal effects were frequently reported in some studies. Common side effects associated with ivermectin included diarrhea, nausea, and dizziness.

Recommendation

Strong recommendation against use of ivermectin to treat COVID-19 (QoE: low).

What is the effect of azithromycin on disease progression in patients with COVID-19 compared to no treatment?

Narrative synthesis of evidence

Azithromycin was assessed in four randomized trials (1 in outpatients and 3 in hospitalized patients). In our analysis, it had no effect on 28-day mortality [RR 1.01; 95% CI 0.92–1.10), risk of disease progression (RR 0.94; 95% CI 0.79–1.14 for mechanical ventilation or ECMO; Table 8), or need for supplemental oxygen [RR 0.84; 95% CI 0.38–1.85). Azithromycin was assessed within the RECOVERY trial which allocated 2582 hospitalized patients to azithromycin and 5181 to SOC; 28-day mortality was similar between groups (RR 0.97, 95% CI 0.87–1.07; p=0.50) [49].

COALITION and COALITION II were open-label randomized trials assessing HCQ, HCQ plus azithromycin, azithromycin, and SOC [20, 50]. The primary endpoint (clinical status at day 15 assessed by 7-grade ordinal scale) was not affected by any of the study drugs in either trial [20, 50]. Azithromycin was not associated with better outcomes in hospitalized patients [51] or outpatients [52].

Safety

Rates of adverse events and severe adverse events were similar in patients receiving azithromycin or SOC [49, 50, 52]. In the only study that assessed azithromycin and HCQ, adverse events and prolongation of the QTc interval were more frequent in patients receiving HCQ or HCQ plus azithromycin compared to controls [20].

Recommendation

Strong recommendation against use of azithromycin for COVID-19 (QoE: high for 28-day mortality, low for disease progression)

What is the effect of colchicine treatment on mortality or disease progression in patients with mild COVID-19 compared with no treatment?

Narrative synthesis of evidence

More than 30 trials have assessed the role of colchicine in COVID-19. Five were considered to define the current position statement. Overall, colchicine had no impact on mortality (RR 1.00; 95% CI 0.93–1.07) or need for mechanical ventilation [RR 1.01; 95% CI 0.91–1.13; Table 9).

COLCORONA compared colchicine to placebo in 4488 outpatients with COVID-19. The primary composite endpoint – death or hospitalization for COVID19 – occurred in 4.7% and 5.8% of patients receiving colchicine and placebo, respectively (OR 0.79; 95% CI 0.61–1.03; p=0.08). Rates of hospitalization and mechanical ventilation and mortality were similar between two groups [53]. Colchicine showed promising results in small preliminary RCTs [54, 55]. However, recent unrefereed results of RECOVERY comparing 28-day mortality in patients receiving colchicine (n=5160) or SOC (n=5730) showed no benefit (RR 1.01; 95% CI 0.93–1.10; p=0.77); this finding was similar in all pre-specified subgroups and in those with SARS-CoV-2 infection confirmed by molecular analysis [56].

Safety

Colchicine has known bone marrow toxicity and several dose-dependent gastrointestinal adverse effects [57]. In COLCORONA, the rate of serious adverse events was 4.9% and 6.3% (p=0.05) and drug-related adverse events were 24.2% and 15.5% (p<0.0001) in the intervention and placebo groups, respectively. Gastrointestinal adverse events were significantly increased with colchicine (23.9% vs 14.8%, p<0.0001) as was diarrhea (13.7% vs 7.3%, p<0.0001) [53]. In the GRECCO trial, no serious adverse events were reported, while adverse events were similar in the two groups with the exception of diarrhea which was mainly seen with colchicine (45.5% vs 18%; p=0.003) [54].

Recommendation

Strong recommendation against use of colchicine for COVID-19 (QoE: high).

What is the effect of corticosteroid treatment on mortality in patients with mild COVID-19 compared with no treatment?

Narrative synthesis of evidence

The evidence involved 5789 patients from 9 RCTs [58-66]. The RR for mortality was significantly lower in patients who received corticosteroids compared to SOC [RR 0.83; 95% CI 0.73–0.99). Corticosteroid treatment was also associated with reduced need for mechanical ventilation [RR 0.88; 95% CI 0.79–0.97; Table 10).

The results of meta-analyses are largely influenced by the RECOVERY trial which enrolled 83% of patients [58]. In RECOVERY, corticosteroid (dexamethasone) provided greater mortality benefits in patients requiring invasive mechanical ventilation (29.3% vs 41.4%) or oxygen support without invasive mechanical ventilation (23.3% vs 26.2%) at randomization [58]. Of the remaining 7 studies (5-11), despite lower mortality with corticosteroid treatment in several trials, some failed to detect significant differences, and some were terminated early based on the results of RECOVERY. In patients who did not require oxygen, corticosteroids likely increased mortality (RR 1.27; 95% CI 1.00–1.61; 1535 patients in one study) and the composite of invasive mechanical ventilation or death [58] (Table 11).

Safety

There was no significant difference between corticosteroid and SOC considering severe adverse events and superinfections. However, corticosteroids are associated with an increase in hyperglycemia. Indirect evidence of corticosteroid use in patients with similar indications has shown no difference in the incidence of gastrointestinal bleeding, superinfections, neuromuscular weakness, or neuropsychiatric effects (Table 10).

Recommendation

Strong recommendation for systemic corticosteroids for treatment of patients with severe and critical COVID-19 (QoE: moderate).

Strong recommendation against the use corticosteroids to treat patients with non-severe COVID-19 (QoE: moderate).

What is the effect of empirical antibiotic treatment on mortality in patients with severe COVID-19 compared to no treatment?

Narrative synthesis of evidence

Several RCTs have not found any effect of azithromycin compared with SOC [49, 50, 52]. In the absence of RCTs assessing antibiotic use in patients with COVID-19 complicated with bacterial coinfections or secondary infections, general principles of antimicrobial stewardship should be applied [67]. Given the low rate of bacterial coinfections, only patients with clinical or radiological suspicion of an associated bacterial infection should receive empirical antibiotics when COVID-19 is diagnosed or when hospitalization is needed.

Recommendation

Insufficient evidence to make a proper recommendation. Antibiotics should not be routinely prescribed in patients with COVID-19 unless bacterial coinfection or secondary infection is suspected or confirmed.

What is the effect of convalescent plasma on mortality in patients with severe COVID-19 compared with no treatment?

Narrative synthesis of evidence

Nine RCTs comparing convalescent plasma with SOC in >12,800 patients with COVID-19 were considered [68-76]. Convalescent plasma did not confer a benefit compared with SOC in 28-day mortality (RR 0.93; 95% CI 0.79–1.10), need for mechanical ventilation (RR 0.98; 95% CI 0.89–1.08), or ICU admission (RR 0.75; 95% CI 0.36–1.59; Table 12). Within RECOVERY, 5795 patients received convalescent plasma and 5763 SOC; 28-day mortality was similar between groups (24% vs. 24%; RR 1.00; 95% CI 0.93–1.07; p=0.95) [68].

PLACID was a multicenter open-label RCT at 39 centers in India enrolling 464 hospitalized adults with moderate-severe COVID-19 [69]. The primary outcome of progression to critical disease or all-cause mortality at 28 days after enrolment was similar between groups (risk difference 0.008; 95% CI -0.062–0.078) (RR 1.04; 95% CI 0.71–1.54).

PlasmAr was a double-blind, placebo-controlled, multicenter trial involving 12 sites in Argentina enrolling patients with severe COVID-19 pneumonia randomized to

receive convalescent plasma (n=228) or placebo (n=105). The primary outcome (clinical status 30 days after intervention) was similar between groups (OR 0.83; 95% CI 0.52–1.35; p=0.46) [76].

Other smaller RCTs found no significant differences in outcomes in patients with moderate-severe [71] or severe-critical COVID-19 [70, 71, 75]. Only one study showed a benefit for convalescent plasma administered in older adult patients within 72 hours after onset of mild COVID-19 symptoms. Progression to severe COVID-19 occurred in 13 of 80 (16%) patients receiving plasma and in 25 of 80 (31%) receiving placebo (RR 0.52; 95% CI 0.29–0.94; p=0.03).

Safety

In general, adverse events were not increased compared to controls [68, 69, 72, 74]. Some studies reported higher rates of serious adverse events [76] or a small number of infusion-related adverse events [73, 75, 76].

Recommendation

Strong recommendation against use of convalescent plasma for COVID-19 (QoE: moderate for mortality, high for mechanical ventilation).

What is the effect of remdesivir on mortality or mechanical ventilation in patients with severe COVID-19 compared to no treatment?

Narrative synthesis of evidence

Our analysis showed that remdesivir probably decreases death slightly in hospitalized patients who do not require ventilation [RR 0.76; 95% CI 0.57–1.02) with uncertain effects on patients undergoing ventilation [RR 1.2; 95% CI 0.98–1.78). Additionally, remdesivir may decrease the need for invasive mechanical ventilation or ECMO (RR 0.57; 95% CI 0.42–0.79; Table 13).

In a double-blind, randomized trial in China enrolling 237 patients with severe COVID-19, time to clinical improvement (hazard ratio (HR) 1.23; 95% CI 0.87–1.75) and mortality rate (14% vs. 13%) were similar with remdesivir and placebo [77]. In SOLIDARITY, 2750 patients were assigned to remdesivir and 2708 to SOC with no difference in 28-day mortality (RR 0.95; 95% CI 0.81–1.11) [12]. ACTT-1 was a multinational, randomized, placebo-controlled trial of remdesivir (given for up to 10 days or until death or discharge) in 1062 patients with confirmed COVID-19. Compared

with placebo, remdesivir resulted in faster time to recovery in the overall population (median 10 vs. 15 days; RR for recovery 1.29; 95% CI 1.12–1.49), but not in the subset on mechanical ventilation or ECMO at baseline (RR for recovery 0.98, 95% CI 0.70–1.36) [78]. Among patients on oxygen supplementation but who did not require high-flow oxygen or ventilatory support (noninvasive or invasive), there was a significant mortality benefit (4.0% vs. 12.7%; HR 0.30; 95% CI 0.14–0.64).

An open-label trial randomized hospitalized patients with moderate COVID-19 pneumonia to a 10-day (n=197) or 5-day course of remdesivir (n=199) or SOC (n=200). A 5-day course (odds ratio (OR) 1.65; 95% CI, 1.09–2.48; p=0.02) of remdesivir, but not a 10-day course (p=0.18), was associated with better clinical status at day 11 vs. SOC. No difference in all-cause 28-day mortality was seen [79]. Another open-label randomized trial compared 5-day to 10-day remdesivir in patients with severe COVID-19. The primary outcome was clinical status at day 14 and was similar between groups (p=0.18) [80]. A small and likely underpowered RCT in India did not show clinical improvement with remdesivir compared to SOC [81]. Lastly, a large observational trial suggested mortality benefit in patients treated with remdesivir compared to those not treated with remdesivir [82].

Safety

Remdesivir was associated with higher rate of adverse events in two studies, especially when administered for 10 days [79, 80]. These included nausea, hypokalemia, headache, and decrease in eGFR. However, a lower rate of serious adverse events was observed in one RCT [77].

Recommendation

Conditional recommendation for use of remdesivir for COVID-19 in hospitalized patinents not requiring mechanical ventilation or ECMO (QoE: moderate).

What is the effect of favipiravir on mortality or mechanical ventilation in patients with mild-moderate COVID-19 compared to no treatment?

Narrative synthesis of evidence

Favipiravir has shown rapid viral clearance and faster clinical improvement of patients with COVID-19 [83]. Certainty of evidence is very low for all-cause mortality, admission to ICU, and need for mechanical ventilation. In recently published RCTs, it

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was found that transfer to ICU, adverse events, and mortality in patients with mild-moderate COVID-19 treated with favipiravir was not significantly different compared with SOC [84]. Several ongoing clinical trials will further substantiate the role of favipiravir [84-86].

Recommendation

Insufficient evidence to make a recommendation.

Is antifungal prophylaxis associated with a lower incidence of coronavirusassociated pulmonary aspergillosis (CAPA) in mechanically-ventilated patients with critical COVID-19 compared to no prophylaxis?

Narrative synthesis of evidence

No antifungal agent is currently approved for prophylaxis in ICU-patients. Recently, posaconazole prophylaxis has been evaluated in ICU patients with severe influenza to prevent influenza-associated pulmonary aspergillosis (IAPA) [87]. Posaconazole was well tolerated and was discontinued prematurely in 9 of 37 patients for causes unrelated to treatment. No cases of IAPA were observed during posaconazole prophylaxis, but the strategy failed as 71% of cases had IAPA on ICU admission that required immediate antifungal therapy [87]. Although CAPA occurs at a median of 7 days after ICU admission and may thus benefit from prophylaxis, there are currently no studies that support this approach in COVID-19 patients in the ICU. Current guidelines and expert guidance do not recommend antifungal prophylaxis in critically-ill COVID-19 patients [88, 89].

Recommendation

Insufficient evidence to make a recommendation.

What is the effect of tocilizumab on mortality or mechanical ventilation in patients with moderate or severe COVID-19 compared to no treatment?

Narrative synthesis of evidence

Tocilizumab has been assessed in 9 RCTs with conflicting results [90-99]. Most of the smaller trials did not show any mortality benefit [90, 93-95, 100, 101]. Conversely, REMAP-CAP and RECOVERY showed small but significant benefit.

REMAP-CAP is an ongoing international, multifactorial, adaptive platform trial including ICU patients randomly assigned to receive tocilizumab, sarilumab, or SOC. The primary outcome was respiratory and cardiovascular organ support-free days. Overall, those with tocilizumab had an in-hospital mortality of 27% compared to 36% for controls, and a median of 10 to 11 organ support-free days compared with 0 days for controls [96].

Within RECOVERY, 4116 patients were assigned to tocilizumab or SOC if they had oxygen saturation <92% on ambient air or required oxygen therapy with evidence of systemic inflammation (C-reactive protein ≥75 mg/l). Overall, 29% patients receiving tocilizumab and 33% receiving SOC died within 28 days (RR 0.86; 95% CI 0.77–0.96; p=0.007) [97].

Tocilizumab is associated with reduced mortality (RR 0.89; 95% CI 0.82–0.98) in 9 RCTs and a lower need for mechanical ventilation (RR 0.81; 95% CI 0.80–0.93) in 4 RCTs (Table 14). One possible explanation for the different results among RCTs is that many were conducted in the early stages of the pandemic before corticosteroids were established as SOC. In a recent systematic review, a clear benefit of combination of IL-6 blockers and corticosteroids was noted [102].

Safety

Tocilizumab likely has little impact on adverse or serious adverse events, septic shock, or clinical progression. The effect of tocilizumab on other outcomes is uncertain.

Recommendation

We recommend use of tocilizumab for treatment of severe COVID-19 (QoE: moderate for mortality, high for mechanical ventilation).

Is intermediate dose of low-molecular weight heparin (LMWH) associated with lower mortality in mechanically-ventilated patients with critical COVID-19 compared to prophylactic dose?

Narrative synthesis of evidence

The use of enoxaparin was assessed in one RCT (INSPIRATION) assessing 562 critically-ill adult patients with COVID-19 admitted to the ICU and followed for 90 days, and randomly allocated to receive intermediate dose or prophylactic dose anticoagulation for 30 days [103]. The primary outcome was a composite including all-

cause mortality, which was similar between groups (HR 1.21; 95% CI 0.95–1.55; p==0.11). RR for all-cause mortality was 1.09 (95% CI 0.78–1.53) (Table 15). In addition, another RCT by the investigators from the REMAP-CAP Platform found clinical benefit from therapeutic dosages of enoxaparin among non-critical COVID-19 patients [104]. However, an analysis restricted to critically-ill patients found no benefit on the primary outcome (ordinal scale combining in-hospital mortality and days free of organ support to day 21) (adjusted OR 0.87, 95% CI 0.70–1.08) [105].

Safety

The main safety outcome in the RCT was major bleeding. There were 7 (2.5%) major bleedings in the intermediate dose group (3 fatal) and 4 (1.4%) major bleedings in the standard-dose group (0 fatal) (HR 1.82; 95% CI 0.53–6.24) [103].

Recommendation

We recommend against the use of intermediate dose of LMWH in critically-ill patients with COVID-19 (QoE: moderate).

We recommend the use of intermediate or therapeutic doses of LMWH in non-criticallyill patients with COVID-19 only in the context of a clinical trial (QoE: moderate).

What is the effect of treatment with interferon β -1a on mortality of critically-ill patients with COVID-19 compared to no treatment?

Narrative synthesis of evidence

Interferon β -1a was not associated with lower 28-day mortality (RR 1.07; 95% CI 0.91–1.27; Table 16). Most patients were enrolled in the SOLIDARITY trial. The primary endpoint was 28-mortality and occurred in 243 of 2050 patients receiving interferon and in 216 of 2050 receiving SOC (RR 1.16; 95% CI 0.96–1.39; p=0.11) [12]. Consistent results were obtained in the subgroup of patients needing mechanical ventilation (RR 1.40; 95% CI 0.82–2.40). A second smaller open-label, single-center study in Iran showed no benefit of interferon β -1a in addition to SOC [106].

In addition to these two trials, another two RCTs are available [107, 108]. Interferon β -1a was not associated with clinical improvement in either trial.

Safety

Some studies documented a higher rate of adverse events in patients treated with interferon β -1a compared with controls [109], whereas others do not [12, 106]. Historically, use of interferons in other settings has been associated with several side effects including thrombotic microangiopathy, hepatic injury, nephrotic syndrome, and depression with suicidal ideation. Interferon β -1a is also associated with immune reactions that can produce flu-like symptoms.

Recommendation

Strong recommendation against use of interferon β -1a in severe COVID-19 patients (QoE: moderate).

Description of the developing group

MB chaired the panel, supervised the work, selected and voted for PICO questions and for other relevant decision, performed literature search and drafted and approved the manuscript.

OA, AB, LB, OE, RK, JRPP, NP, MS, BGZ, ST, PEV, IZS selected and voted for PICO questions and for other relevant decision, performed literature search and drafted and approved the manuscript.

JRB supervised the work of the panel, selected and voted for PICO questions and for other relevant decision, performed literature search and drafted and approved the manuscript.

Conflict of interest

MB, OA, AB, LB, OE, RK, X, JR-PP, NRP, BGSZ, MS, TS, IZS JR-B: no conflicts to declare.

PEV: reports research grants from Gilead Sciences, MSD, Pfizer and F2G; he is a speaker for Gilead Sciences, Mundipharma F2G, and MSD; and is on the advisory boards for Pfizer, MundiPharma, Cidara, MSD, and F2G.

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Updating

The panel will meet monthly to assess the need for further update of the present document. Our goal will be an optimisation of the guideline development process to allow update of individual recommendations as soon as new evidence becomes available. More specifically, during the monthly meeting the panel will propose new PICOs or revision of prior PICOs. The methodology will be the same of the present document.

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Table 1. ADOLOPMENT criteria used to determine the suitability of the existing evidence synthesis (need for updating the literature search and for revising the grading of the quality of the evidence).

Criterion	New systematic review (a systematic review that does not qualify as major or minor update)				Major update (first criterion applies and any of the following)	Minor update (all must apply	
Prior review (for	No credible available system	matic r	eview e	exists	A credible systematic	A credible	systematic
question)	for the question*				review exists*	review exists*	
Full text reviewed for the question of interest	N/A				>20	≤20	
New studies	N/A				>5	<u><</u> 5	
Evidence profile available	N/A				Not available	Available	
Outcomes all	Not all important outcomes			Ó	All important outcomes	All important	outcomes
addressed	addressed				addressed	addressed	
Type of studies	Search for observational studies						

^{*}A credible available review is one that has publicly available data, has been conducted in the past 4 months (or a different timescale if deemed appropriate by the drafting group), scores highly on the AMSTAR or another tool, has a reproducible search strategy, meta-analysis (that can be reproduced), existing accessible risk of bias evaluation of individual studies (that can be reproduced).

Table 2. Summary of reccomendations and dosages

Severity of	Treatment	Dosages	European	Comments
disease/setti	recommended		Medicine	
ng			Agency	
			Authorizati	
			on *	
Mild COVID-	AntiSpike	Bamlanivi	Rolling	Only in
19	monoclonal	mab 700	review	patients with
Outpatient	antibodies	mg +		risk factors for
setting	(conditional	etesemiva		disease
	recommendation)	b 1400 mg	Ç.	progression#
		 Casirivima 		
		b 1200mg		
		+		
		Imdevimab		
		1200 mg		
Mild COVID-	Casirivimab/imdevi	Casirivimab 4 g	Rolling	
19	mab	plus imdevimab 4 g	review	
Inpatient	(conditional			
setting	recommendation)			
	Remdesivir	200 mg IV loading	Approved	
	(conditional	dose, followed by		
	recommendation)	100 mg daily for 5		
		days		
Severe or	Casirivimab/imdevi	Casirivimab 4 g	Rolling	
Critical	mab	plus imdevimab 4 g	review	
COVID-19	(conditional			
	recommendation)			
	Dexamethasone	6 mg PO or IV daily	Approved	recommended
	(strong	for 10 days or until		in patients
	recommendation)	discharge		receiving
				oxygen
				supplement
	Tocilizumab	8 mg per kg of	Approved	
	(Strong	actual body weight		
	recommendation)	(up to a maximum		
		of 800 mg), as an		
		intravenous		
		infusion over a		

	period of 1 hour. A		
	second dose may		
	be repeated 12 to		
	24 hours later		
Remdesivir	200 mg IV loading	Approved	Not
(conditional	dose, followed by		recommended
recommendation)	100 mg daily for 5		in patients
	days		requiring high-
			flow oxygen
			supplementati
		C	on

^{*}https://www.ema.europa.eu/en/human-regulatory/overview/public-health-threats/coronavirus-disease-covid-19/treatments-vaccines/covid-19-treatments accessed October 20th 2021.

- # Risk factors for disease progression to consider for mAb treatment in adult patients:
 - Body mass index ≥ 35
 - Chronic kidney disease
 - Diabetes
 - Immunosupressive disease
 - Age ≥ 65 years
 - Age ≥ 55 years and at least one of the following:
 - o Cardiovascular disease
 - Hypertension
 - Chronic obstructive pulmonary disease or other chronic respiratory conditions

Table 3. Grade evidence profile PICO1: Hydroxychloroquine for COVID-19

Hydroxychloroquine for COVID-19
People: Patients with COVID-19
Settings: Inpatients (15 studies) Outpatient (5 studies) Intervention: Hydroxychloroquine

Comparison: No trea Outcomes	Absolute	Effect	Relative	Numbe	Certainty of
	Without Hydroxychloroquine	With Hydroxychloroquin e	effect (95% CI)	r of studies	the evidence (GRADE)
All-cause mortality	168 178 per 1000 Difference: 10 more per 1000 (95% CI: -5 to 27)		RR 1.06 (0.97to 1.16)	19 [12- 16, 18- 21, 23, 25-28, 110- 114]	⊕⊕⊕ ⊕ High
				(10,382 patients	
Invasive mechanical ventilation or ECMO	85 per 1000 Difference: 7 m (95% Cl: -		RR 1.08 (0.91 to 1.28)	8 [14, 15, 18- 20, 28, 112, 114] (5701 patients	⊕⊕⊕ ⊕ High
Hospitalization (end of follow-up)	55 per 1000 Difference: 18 f e (95% CI: -		RR 0.68 (0.41to 1.13)	5 [26- 28, 110, 113] (1345 patients	Low (serious imprecision and serious risk of bias)
Clinical deterioration (within 28 days of treatment begin)	89 per 1000 Difference: 17 fo (95% CI: -5	58 to 79)	RR 0.81 (0.35 to 1.89)	1 [19] (247 patients)	Low (very serious imprecision)
	756 per 1000	794 per 1000	RR 1.05 (0.91 to 1.2)	1 [19]	

Clinical Improvement (within 28 days of treatment begin)				(247 patients)	Low (very serious imprecision)
Discharge for hospital (within 28 days of treatment begin)		680 per 1000 fewer per 1000 -28 to 7	RR 0.98 (0.96 to 1.01)	5 [12, 19, 111, 112, 114] (7365 patients	⊕⊕⊕ ⊕ High
Adverse events (end of follow-up)		538 per 1000 6 more per 1000 68 to 419)	RR 1.67 (1.21 to 2.3)	11 [13, 14, 19- 23, 27, 28, 110, 115] (2077 patients	Moderate (serious risk of bias)
Serious adverse events (end of follow-up)		74 per 1000 more per 1000 -10 to 25)	RR 1.09 (0.86 to 1.37)	11 [13, 19, 20, 22, 23, 25, 27- 29, 113, 115] (2721 patients	Moderate (Serious risk of bias)
	idence interval; RR: Risk 6, 18-29, 110-115].	ratio		,	
	dopted: Australian p.org/#/guideline/5446/section	National <u>on/78675</u>).	COVID-19	Evidence	Taskforce

Table 4. Grade evidence profile PICO2: Bamlanivimab for COVID-19

People: Patients with COVID-19

Settings: Outpatients

Intervention: bamlanivimabComparison: No treatment

Outcomes	Absolute	Effect*	Relative	Numbe	Certainty of
	With bamlanivimab	Without bamlanivimab	effect (95% CI)	r of studies	the evidence (GRADE)
Hospitalization (within 29 days from	5/309 (1.6%)	9/143 (6.3%)	RR 0.26 (0.09 to 0.75)	1 [30]	$\oplus \oplus \ominus$
treatment)	Difference: 47 f (95% CI:-5	-		(452 patients	Low (very serious imprecision)
Serious adverse events (end of follow-up)	0/309 (0%) per 1000 Difference: 6 fe (95% CI: -	•	RR 0.15 (0.01 to 3.78)	1 [30] (452 patient)	Low (very serious imprecision)

95% CI: 95% Confidence interval; RR: Risk ratio

References: [30].

Evidence adopted: Infectious Disease Society of America (IDSA) guidelines available at https://www.idsociety.org/practice-guideline/COVID-19-guideline-treatment-and-management/.

Table 5. Grade evidence profile PICO2: Bamlanivimab in combination with etesevimab for COVID-19

People: Patients with COVID-19

Settings: Outpatients

Intervention: bamlanivimab/etesevimab

Comparison: No treatment

Outcomes	Absolute	Relative	Numbe	Certainty of	
	With bamlanivimab/	Without	effect	r of	the
	etesevimab	bamlanivimab/	(95% CI)	studies	evidence
		etesevimab			(GRADE)
All-cause mortality (within 29 days from	0/518 (0%)	10/517 (1.9%)	RR 0.05 (0.00 to 0.80)	1 [116]	$\oplus \oplus$
treatment)	Difference: 19 fe		C	(1035	$\Theta\Theta$
	(95% CI: -3	31 to -7)		patients	Low
				, pationto	(due to
)	serius
					imprecisio
					n)
Hospitalization (within 29 days from	11/518 (2.1%)	36/517 (7.0%)	RR 0.30 (0.16 to 0.59)	1 [116]	$\oplus \oplus$
treatment)	Difference: 49 fe		1	(1035	$\Theta\Theta$
	(95% CI:-58		patients	Low	
				'	(due to
)	serius
					imprecisio
Serious adverse	7/518 (1.4%)	E/E47 (40/)	RR 1.40	1 [116]	n)
events	7/510 (1.4%)	5/517 (1%)	(0.45 to 4.37)	(1035	$\oplus \oplus \ominus$
(end of follow-up)	Difference: 4 more per 1000		† `	patients	Θ
	(95% CI: -5	to +33))	Low
					(serious
					imprecision)

95% CI: 95% Confidence interval; RR: Risk ratio

References: [116].

Evidence adopted: Infectious Disease Society of America (IDSA) guidelines available at https://www.idsociety.org/practice-guideline/COVID-19-guideline-treatment-and-management/.

Table 6. Grade evidence profile PICO3: Casirivimab combined with imdevimab for COVID-19

People: Patients with COVID-19

Settings: Outpatients

Intervention: casirivimab (1200 mg) combined with imdevimab (1200 mg)

Comparison: No treatment

Outcomes	Absolute	Effect	Relative	Numbe	Certainty of
	With casirivimab combined with imdevimab	Without casirivimab combined with imdevimab	effect (95% CI)	r of studies	the evidence (GRADE)
All-cause mortality (within 29 days from	1/736 (0.1%)	1/748 (0.4%)	RR 1.02 (0.06 to	1 [35]	$\oplus \oplus$
treatment)	Difference: 0 fe (95% Cl: -		16.20)	(1484 patients	Low due to very serious imprecision)
Hospitalization (within 29 days from treatment)	6/736 (1.9%) Difference: 22 fe (95% CI:-2		RR 0.27 (0.11 to 0.65)	1 [35] (1484 patients	Moderate (Due to serious imprecision)
Serious adverse events (end of follow-up)	50/3688 (1.2%) Difference: 27 fe (95% CI: -3	-	RR 0.34 (0.24 to 0.48)	1 [35] (5531 patients)	Moderate (Due to serious imprecision

95% CI: 95% Confidence interval; RR: Risk ratio

References: [35].

Evidence adopted: Infectious Disease Society of America (IDSA) guidelines available at https://www.idsociety.org/practice-guideline/COVID-19-guideline-treatment-and-management/.

Table 7. GRADE evidence profile for PICO 4: Ivermectin for COVID-19.

Ivermectin vs Standard care

People: Adult patients with COVID-19

Setting: Inpatients (10 studies), Outpatients (7 studies)

Intervention: Ivermectin

Comparison: Standard Care (15 studies), HCQ (1 study), Lopinavir/ritonavir (1 study)

Outcomes	Absolute	Effect	Relative	Number	Certainty of
	Without Ivermectin (Standard Care)	With Ivermectin	_ effect (95% CI)	of studies	the evidence (GRADE)
All-cause mortality Within 28 days of commencing treatment	53 per 1000 Difference: 31 fe (95% CI: 43 fewe		RR 0.41 (0.19 to 0.92)	6 [17, 45, 47, 117- 119] (1079 patients)	Low (serious risk of bias and serious imprecision
Mechanical ventilation Within 28 days of commencing treatment	40 per 1000 Difference: 4 fer (95% CI: 31 fewe		RR 0.75 (0.23 to 2.43)	4 [88, 117, 118] (497 patients)	Low (very serious imprecsion)
Serious adverse events End of treatment	7 8 per 1000 Difference: 25 more per 1000 (95% CI: 19 fewer to 89 more)		RR 1.12 (0.21 to 5.88)	6 [42-44, 47, 120, 121] (644 patients)	⊕⊕⊖⊖ Low (Very Serious imprecision,)
Adverse events End of treatment	497 per 1000 Difference: 25 fe (95% Cl: 70 fewe		RR 0.95 (0.86 to 1.05)	7 [42-44, 47, 120, 121]	⊕⊕⊖⊖ Low (Serious imprecision,

				(805 patients)	serious risk of bias)
ICU admission End of follow-up	115 per 1000	61 per 1000	RR 0.53 (0.11 to 2.51)	2 [44, 45]	⊕⊕⊖⊖ Low
	Difference: 54 fev (95% CI: 102 fewer			(143 patients)	(Serious imprecision, serious risk of bias)
Discharge from hospital	868 per 1000	920 per 1000	RR 1.06 (0.99 to 1.12)	4 [17, 43, 118,	⊕⊕⊖⊖ Low
Within 28 days of commencing treatment	Difference: 52 mg (95% CI: 9 fewer to		Š	122] (342 patients)	(Serious imprecision, serious risk of bias)

95% CI: 95% Confidence interval; RR: Risk ratio

References: [17, 39, 40, 42-47, 117-123].

Evidence adopted Australian guidelines for the clinical care of people with COVID-19, Available at: https://app.magicapp.org/#/guideline/5446/section/78706

Table 8. Grade evidence profile PICO5: Azithromycin for COVID-19

Azithromycin vs Standard care

People:Adult patients with COVID-19 (pregnant patients excluded)

Setting: hospital (4 studies), outpatients (1 study) [124], 3 Countries (Iran, Brazil, UK)

Intervention: Azithromycin (500 mg o.d.), 3 to 10 days.

Comparison: Standard Care

Patients in both intervention and comparator arms also receiving HCQ in 2 studies [20, 50] and HCQ+ LPV/r in 1

study [51].

Outcomes	Absolute	Effect*	Relative effect	Number of	Certainty of the
	Without Azithromycin (Standard Care)	With Azithromycin	(95% CI)	studies	evidence (GRADE) [†]
All-cause mortality Within 28 days of commencing treatment	172 per 1000 Difference: 2 m (95% Cl: 14 fewe		RR 1.01 (0.92 to 1.10)	4 [50, 51, 124, 125] (9595 patients)	⊕⊕⊕ High
Supplemental oxygen	24 per 1000	20 per 1000	RR 0.84 (0.38 to 1.85)	1 [124]	⊕⊕⊖⊖ Low
Within 28 days of commencing treatment	Difference: 4 fe (95% CI: 15 fewe			(1122 patients)	(Very serious imprecision; only data from one study, due to few events)
Clinical recovery	658 per 1000	632 per 1000	RR 0.96 (0.88 to 1.05)	1 [124]	⊕⊕⊖⊖ Low
Within 28 days of commencing treatment	Difference: 26 fe (95% Cl: 79 fewe			(1129 patients)	(Very serious imprecision; wide confidence intervals, only data from one study)
	60 per 1000	56 per 1000	RR 0.94 (0.79 to 1.14)		

Ba - h - · · · ·	<u> </u>		<u> </u>	0.540.4	0.00-
Mechanical				2 [124,	$\oplus \oplus \oplus \oplus$
ventilation or				125]	High
ECMO	Difference: 4 few (95% CI: 13 fewer		(8433		
Within 28 days of	(93% Cl. 13 lewel		,		
commencing				patients)	
treatment					
Serious	194	219	RR 1.13	2 [20,	$\oplus \oplus \oplus \ominus$
adverse events	per 1000	per 1000	(0.90 to 1.42)	50]	Moderate
End of treatment				(877	(serious
	Difference: 25 mc			(011	imprecision;
	(95% CI: 19 fewer	to 89 more)	Ç.	patients)	wide confidence
					intervals)
Adverse events	337	394	RR 1.17 (0.91 to 1.50)	1 [20]	0000
End of treatment	per 1000	per 1000	(0.91 to 1.50)	(438	Low
	I	(0)		(430	(Very serious
				patients)	imprecision;
	Difference: 57 mc (95% CI: 30 fewer t			wide confidence	
	(95% CI. 30 lewer t			intervals, only data from one	
					study)
ICU admission	18 per 1000	9 per 1000	RR 0.48 (0.17 to 1.35)	2 [124]	$\oplus \oplus \ominus \ominus$
End of follow-up	ps. 1000	poi 1000	(0117 10 1100)	(1231	Low
				((Very serious imprecision,
	Difference: 9 few (95% CI: 15 fewer			patients)	due to few
	(con on io ionion	10 0			everns)
Discharge from	586	539	RR 0.92	2 [50,	$\oplus \oplus \oplus \ominus$
hospital	per 1000	per 1000	(0.72 to 1.19)	2 [30, 125]	Moderate
Within 28 days of				120]	Moderate
commencing treatment				(8162	(serious
	Difference: 47 fev (95% CI: 170 fewer				imprecision;
	(**************************************	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,		patients)	wide confidence
					intervals)
Duration of			-	2 [20,	$\oplus \oplus \ominus \ominus$
hospital stay Mean				51]	Low
	Difference: 0.41 lower (MD) (95% CI: 2.42lowerto 1.59 higher)			(442	(serious
	(90% OI. 2.42IUWERK	, i.əə iligilet)		(442	inconsistency and
				patients)	imprecision; wide confidence
					intervals)
	13	12		1 [125]	
ĺ					

Duration of hospital stay	-	(7764	$\oplus \oplus \oplus \ominus$
		patients)	Moderate
Median			
			(serious
			imprecision;
			only data from1
			study)

95% CI: 95% Confidence interval; RR: Risk ratio

References: [20, 50, 51, 124, 125].

Evidence adopted Australian guidelines for the clinical care of people with COVID-19, Available at: https://app.magicapp.org/#/guideline/5446/section/78706

Table 9. Grade evidence profile PICO6: Colchicine for COVID-19

People:Adult patients with COVID-19 (pregnant patients excluded)

Setting: Hospital

Intervention: Colchicine **Comparison:** Standard Care

Outcomes	Absolute	Effect	Relative effect	Number of	Certainty of the
	Without Colchicine (Standard Care)	With Colchicine	(95% CI)	studies	evidence (GRADE)
All cause mortality within 21-28 days	149 per 1000	149 per 1000	RR 1.00 (0.93 - 1.07)	4 [53-55, 126]	⊕⊕⊕⊕ High
of treatment administration	0 fewer pe (CI 95% 10 fewe			(15968 patients)	
Disease progression Increase of 2 grades on 7-grade scale; 21 days after commencing treatment	140 per 1000 18 per 1000 Difference: 4 122 fewer per 1000 (95% CI: 187 fewer to 3 more)		RR 0.13 (0.02 - 1.02)	1 [54] (105 patients)	Uvery serious imprecision; only data from one study, due to few events)
Invasive mechanical	80 per 1000	81 per 1000	RR 1.01 (0.91 - 1.13)	3 [53, 54, 126]	⊕⊕⊕⊕ High
within 21-28 days of treatment administration	Differen		·	(15404 patients)	
Serious adverse events End of treatment	61 per 1000 Difference: 13 m (95% CI: 24 fewe		RR 0.78 (0.61 to 1.00)	2 [53, 54] (4517 patients)	⊕⊕⊕⊖ Moderate (serious imprecision; wide confidence intervals)
	158	305	RR 1.93		

Adverse events	per 1000	per 1000	(1.18 to 3.16)	2 [53,	$\oplus \oplus \oplus \ominus$
End of treatment			-	54]	Moderate
	Difference: 147 m (95% CI: 28 more			(4517 patients)	(serious imprecision; wide confidence intervals)

95% CI: 95% Confidence interval; RR: Risk ratio

References: [53-55, 126, 127].

Evidence adopted Australian guidelines for the clinical care of people with COVID-19, Available at: https://app.magicapp.org/#/guideline/5446/section/78673

Table 10. Grade evidence profile of PICO 7: corticosteroids for adult patients with COIVD-19 requiring oxygen supplement

Corticosteroids for severe COVID-19 i.e. patients requiring oxygen including mechanically ventilated patients

People: Patients with COVID-19

Settings: Error! Bookmark not defined.

Intervention: Corticosteroids **Comparison:** No treatment

Comparison: No treatment								
Outcomes	Absolu	ite Effect	Relative	Numbe	Certainty of			
	Without	With	effect	r of	the			
	Corticosteroids	Corticosteroids	(95% CI)	studies	evidence			
		33111333131313			(GRADE)			
All-cause mortality (adults requiring	316	265	RR 0.84	9 [58-	$\oplus \oplus$			
oxygen)	per 1000	per 1000 fewer per 1000	(0.73 to 0.98)	63, 65,				
		ewer – 6 fewer)		66, 128,	$\oplus \ominus$			
	,				Moderat			
				129]	е			
				(5789	(due to serious			
				(3789	inconsisten			
				patients)	cy)			
					,,			
Invasive mechanical	320	282	RR 0.88	1 [58]	$\oplus \oplus$			
ventilation or death (adults requiring	per 1000	per 1000	(0.79 to 0.97)					
oxygen)		s fewer per 1000 ewer – 10 fewer)		(3883	$\oplus \ominus$			
		,		patients	Moderat			
)	е			
				,	due to			
					serious			
					inconsisten			
					су			
Serious adverse events (adults	234	187	RR 0.80	6 [59,	$\oplus \oplus \oplus$			
requiring oxygen)	per 1000	per 1000 ' more per 1000	(0.53 to 1.19)	60, 62,				
		ewer – 44 more)		63, 65,	\bigcup			
	(00)0000	,		128] (696	Moderate			
				patients	(due to serious			
)	inconsistency)			
Superinfection	186	188	RR 1.01	32 [129]	\triangle			
(end of treatment)	per 1000	per 1000	(0.90 to 1.13)	' '	$\Theta\Theta$			
		more per 1000		(6027	$\Theta\Theta$			
	(95% Ci: 19 to	ewer – 24 more)		patients	Low			
				patients	(Due to			
)	serious indirectness			
					and			
Hyperglycemia	200	222			imprecision)			
туретујусенна	286	332						

(end of treatment)	Per 1000 Per 1000 Difference: 46 more per 1000 (95% CI 23 more -72 more)		RR 1.16 (1.08-1.25)	24 [129]	Moderate (due to serious indirectness)
Discharge from hospital (within 28 days of treatment begin, adults requiring oxygen)	582 per 1000 Difference: 58 (95% CI: 35 mc	640 per 1000 more per 1000 ore to 87 more)	RR 1.10 (1.06 to 1.15)	2 [58, 66] (4952 patients	Moderat e (due to serious inconsistenc

95% CI: 95% Confidence interval; RR: Risk ratio

References: [58-63, 65, 66, 128, 129].

Evidence adopted: Australian guidelines for the clinical care of people with COVID-19, Available at: https://app.magicapp.org/#/guideline/5477/section/80465

Table 11. GRADE evidence profile PICO7: Corticosteroid for COVID-19 in the subgroup of hospitalized patients not requiring supplemental oxygen

Corticosteroids for mild COVID-19 i.e. patients not requiring oxygen

People: Patients with COVID-19

Settings: Error! Bookmark not defined. Inpatients

Intervention: Corticosteroids **Comparison:** No treatment

Outcomes	Absolute	Effect	Relative	Numbe	Certainty of
	Without	With	effect	r of	the
	Corticosteroids	Corticosteroids	(95% CI)	studies	evidence (GRADE)
All-cause mortality	140 per 1000	178 per 1000	RR 1.27 (1.00 to 1.61)	1 [58]	$\oplus \oplus$
	Difference: 38 m			(1535	$\oplus\ominus$
	(95% CI: 0 more	e to85 more)		patients	Moderat
)	e (serious
		40i			imprecision)
Invasive mechanical ventilation or death	155 per 1000	194 per 1000	RR 1.25 (1.0 to 1.57	1 [58]	$\oplus \oplus$
	Difference: 39 m (95% CI: 0 more			(1535	$\oplus \ominus$
	(00)	,		patients	Moderat
)	e (serious
					imprecision)
Discharge for hospital	804 per 1000	772 per 1000	RR 0.96 (0.9 to 1.01)	1 [58]	$\oplus \oplus$
(within 28 days of treatment begin)	Difference: 32 fe	ewer per 1000		(1535	$\oplus \ominus$
σ,	(95% CI: 80 fewer	er to 8 more)		patients	Moderat
)	e (serious
					imprecision)

95% CI: 95% Confidence interval; RR: Risk ratio

References: [58].

Evidence adopted: Australian guidelines for the clinical care of people with COVID-19, Available at: https://app.magicapp.org/#/guideline/5477/section/80465

Table 12. GRADE evidence profile for PICO 9: convalescent plasma for COVID-19

People:Adult patients with COVID-19 (pregnant patients excluded) **Setting:** hospitalized patients (8 studies), outpatients (1 study)

Intervention: Convalescent plasma

Comparison: Standard Care

Comparison: Standard Care								
Outcomes	Absolute	Effect	Relative	Number	Certainty of			
		1000	effect	of	the evidence			
	Without Convalescent Plasma	With Convalescent Plasma	(95% CI)	studies	(GRADE)			
	(Standard Care)		K					
All-cause mortality	235 per 1000	219 per 1000	RR 0.93 (0.79 to 1.10)	9 [69-72, 74-76,	⊕⊕⊕⊖ Moderate			
Within 28 days of commencing treatment				130, 131].	(due to serious imprecision)			
	Difference: 16 f (95% Cl: 49 few			(12872				
	(00/0 0:: 00/00:			patients)				
Invasive	124 per 1000	122 per 1000	RR 0.98 (0.89 to 1.08)	4 [69, 74,	$\Theta \oplus \Theta \oplus \Theta$			
mechanical	per 1000	per 1000	(0.03 to 1.00)	76, 130]	High			
ventilation								
Within 28 days of commencing	Difference: 2 fe (95% CI: 14 few			(11898				
treatment				patients)				
Serious	176 per 1000	218 per 1000	RR 1.24 (0.81 to 1.90)	2 [71, 76]	000			
Adverse events	po. 1000	J PO. 1000	(0.01.00)		Low			
Within 28 days of		44		(44.4	(Very serious imprecision; wide			
commencing	Difference: 42 r (95% CI: 33 fewe			(414	confidence			
treatment	•			patients)	intervals, only data from one study)			
Adverse events	537	789	RR 1.47	2 [70, 76]	0000			
	per 1000 Difference: 252	per 1000	(0.38 to 5.74)		Low			
	(95% Cl: 333 fewe				(risk of bias; serious			

Within 28 days of commencing treatment				(370 patients)	imprecision; wide confidence intervals, only data from 2 study)
ICU admission	373 per 1000	280 per 1000	RR 0.75 (0.36 to 1.59)	2 [74, 76] (493	⊕⊕⊖⊖ Low
End of follow-up	Difference: 93 fewer per 1000 (95% CI: 239 fewer to 220 more)			patients)	(Risk of bias serious imprecision, due to few events)
Clinical deterioration (progression to	74 per 1000	53 per 1000	RR 0.71 (0.18 to 2.78)	2 [69, 71] (545	⊕⊕⊖⊖ Low
Severe/critical) Within 28 days of commencing	Difference: 21 fewer per 1000 (95% CI: 61 fewer to 132 more)			patients)	(Risk of bias serious imprecision, wide
treatment		2,0			confidence intervals)

95% CI: 95% Confidence interval; RR: Risk ratio

References: [69-72, 74-76, 130, 131].

Evidence adopted: Australian guidelines for the clinical care of people with COVID-19, Available at: https://app.magicapp.org/#/guideline/5477/section/80436

Table 13. GRADE evidence profile for PICO 10: remdesivir for severe COVID-19

Remdesivir for COVID-19

People: Patients with severe COVID-19

Settings: Inpatients

Intervention: Error! Bookmark not defined. Comparison: Error! Bookmark not defined.

Outcomes	Absolute	e Effect	Relative	Numbe	Certainty of
	Without	With	effect	r of	the
	Remdesivir	Remdesivir	(95% CI)	studies	evidence (GRADE)
All-cause mortality	90	68	RR 0.76	5[12,	$\oplus \oplus \oplus$
(hospital, no ventilation)	per 1000	per 1000	(0.57 to 1.02)	77-81]	ΨΨΨ
,	Difference: 22 for (95% CI: 39 few				Θ
	((6,400	Moderat
				patients	е
				pationto ,	(Due to
)	serious imprecision)
All-cause mortality (ventilation)	248	298	RR 1.2	3 [12,	$\oplus \oplus \oplus$
(veritilation)	per 1000 Difference: 50 r	per 1000	(0.98 to 1.47)	77, 78]	9
	(95% Cl: 5 fewe				\Box
				(1,004	Moderat
				patients	е
)	(Due to serious imprecision)
Respiratory failure or ARDS	143	113 per 1000	RR 0.79	2 [77,	$\oplus \oplus$
	per 1000 Difference: 30 f	(0.35 to 1.78)	78]		
	(95% CI: 93 fewer to 112 more)				
	,	,		(1,296	Low (Due to serious
				patients	risk of bias and
				,	serious inconsistency)
					inconcionation by)
Invasive mechanical	225	128	RR 0.57	1	ФФ
ventilation or ECMO (within 28 days of	per 1000	per 1000	(0.42 to 0.79)	[78]	$\oplus \oplus$
treatment start)	Difference: 97 for			[, 0]	$\Theta\Theta$
	(95% CI: 131 few	er-to 47 fewer)		(766	Low
				patients	(Due to serious
)	risk of bias and serious
					inconsistency)
Patients requiring	114	119	RR 1.04	2 [77,	$\oplus \oplus \oplus$
ventilation (within 28 days of treatment	per 1000	per 1000	(0.89 to 1.21)	81]	
start)	Difference: 5 more per 1000 (95% CI: 13 fewer – 24 more)		1		\cup

			1	/= aa 4	
				(5,034	Moderate
				patients	(Due to serious
				·)	imprecision)
				,	
Serious adverse	253	190	RR 0.75	3 [77-	$\Delta \Delta \Delta$
events	per 1000	Per 1000	(0.63-0.89)	79]	$\oplus \oplus \oplus$
End of follow-up	Difference: 63	fewer per 1000	(11111)	(1,865	Θ
	(95% CI 94 fev	wer -28 fewer)		patients	Moderate
	,	,		')	
				,	(Due to
					serious risk of
					bias)
Adverse events	548	570	RR 1.04	3 [77-	$\oplus\oplus\ominus$
End of follow-up	per 1000	per 1000	(0.89-1.21)	79]	$\Phi\Phi$
·	Difference: 22			(1,880	Θ
	(95% CI 60 fev	ver -115 more)		patients	Low
			C)	(due to
					serious risk of
					bias and
					inconsistency)

95% CI: 95% Confidence interval; RR: Risk ratio

References: [12, 77-81].

Evidence adopted: Australian guidelines for the clinical care of people with COVID-19, Available at:

https://app.magicapp.org/#/guideline/5446/section/78660

Table 14. GRADE evidence profile for PICO 13: Tocilizumab for moderate or severe COVID-19.

People: Patients with COVID-19 **Settings:** hospitalized patients **Intervention:** Tocilizumab

Comparison: standard treatment without Tocilizumab

Outcomes	Absolute		Relative	Numbe	Certainty of
	Without Tocilizumab	With Tocilizumab	effect (95% CI)	r of studies	the evidence (GRADE)
All-cause mortality Day 21-28 after treatment start	290 per 1000 Difference: 32 fe (CI 95% 52 fewe		RR 0.89 (0.82 — 0.98)	8 [90- 99, 101] (6481 patients	Moderate ⊕⊕⊕○ (Due to serious imprecision)
Invasive mechanical ventilation or ECMO End-of-follow-up	159 per 1000 Difference: 30 fe (95% CI: 48 fewe		RR 0.81 (0.70 to 0.93)	3 [95, 97, 101] (4248 patients)	⊕⊕ ⊕⊕ High
Admission to ICU End-of-follow-up	423 per 1000 Differece: 76 fe (95% CI: 195 fev		RR 0.82 (0.54-1.23)	4[90, 93, 101] (699 patients	Moderate ①①①① (Due to serious imprecision)
Serious adverse events End-of-follow-up	162 Per 1000 Differece: 18 fe (95% CI: 41 fe		RR 0.89 (0.75 — 1.05)	7 [90, 92, 94- 96, 98, 101] (2309 patients	Moderate ⊕⊕⊕○ (Due to serious imprecision)
Adverse events End-of-follow-up	466 Per 1000 Difference: 28 n (95% CI: 65 few		RR 1.06 (0.86 — 1.3)	6 [90, 92, 94, 95, 98, 101] (1562 patients	Moderate ⊕⊕⊕○ (Due to serious imprecision)

References: [90-99, 101].

Evidence adopted: Australian guidelines for the clinical care of people with COVID-19, Available at: https://app.magicapp.org/#/guideline/5446/section/78668



Table 15. GRADE evidence profile for PICO 14: Low molecular weight heparin for critical COVID-19

Patients or population: mechanical ventilated patients with critical COVID-19

Settings: inpatients

Intervention: intermediate dose of enoxaparin

Comparison: prophylactic dose LMWH

Outcomes	Absolute	Effect	Relative	Numbe	Certainty of
	Risk with prophylactic dose	Risk with intermediate dose	effect (95% CI)	r of studies	the evidence (GRADE) [†]
All-cause mortality follow up: mean 30	409 per 1,000	429 per 1,000	OR 1.09 (0.78 - 1.53)	1 [103]	$\oplus \oplus$
days	Difference: 20 more (CI 95% 58 fewer -105 m			(562	$\oplus \ominus$
	,			patients	Moderat
)	е
				,	(Due to
					serious
					imprecisio n 1 RCT)
Pulmonary embolism	17 per 1000	13 per 1000	OR 0.41 (0.08 - 2.13)	1 [103]	$\oplus \oplus$
	Difference: 10 fe		1	(562	$\Theta\Theta$
	(Margin of error: 16	fewer to 19 more)		patients	Low (serious
)	risk of bias,
					serious imprecision
Major Bleeding	14 per 1000	19 per 1000	OR 1.83 (0.53 - 5.93)	1 [103]	$\oplus \ominus$
	Difference: 11	1	(562	$\Theta\Theta$	
	(Margin of error: 7		patients	Very low	
	7)	

95% CI: 95% Confidence interval; RR: Risk ratio

References: [103].

Evidence adopted:

https://www.hematology.org/-/media/hematology/files/clinicians/guidelines/vte/etd-ash-COVID-19-guideline-recommendation-1a.pdf.

Table 16. GRADE evidence profile for PICO 15: Interferon β -1a for critical COVID-19.

People: Adult patients with COVID-19 (pregnant patients excluded)

Setting: hospitalized (2 studies) patients

Intervention: Interferon β -1a 44 μ g three times per week

Comparison: Standard Care

Outcomes	Absolute Effect		Relative	Number	Certainty of the
	Without Interferon β-1a (Standard Care)	With Interferon β- 1a	effect (95% CI)	of studies	evidence (GRADE)
All-cause mortality	112 per 1000	120 per 1000	RR 1.07 (0.91- 1.27)	2 [12, 106]	⊕⊕⊕⊕ High
Within 28 days of commencing treatment	Difference: 8 n (95% Cl: 10 few		(4181 patients)	Moderate for critical ill (due to serious indirectness)	
Supplemental Ventialtion Within 28 days of commencing treatment	116 per 1000 Difference: 1 fe (95% CI: 20 few	RR 0.99 (0.83 - 1.17)	2 [12, 106] (3912 patients)	Uvery serious imprecision; only data from one study, due to few events)	
Duration of hospital stay Mean days to discharge	12.3 (mean) 14.8 (mean) Difference: 2.55 days higer (95% CI: 0.92 lower to 6.02 higher)			1 [106] (81 patients)	⊕⊕⊕⊕ Very Low (Very serious risk of bias ;very serious imprecision and wide confidence intervals, only data from one study)
Serious adverse events End of follow-up	385 per 1000 Difference: 158 (95% CI: 35 fewe		RR 1.41 (1.09 -1.81)	1 [107] (292 patients)	⊕⊖⊖ Very Low (serious imprecision; serious indirectness)

Adverse events End of follow-up	709 per 1000	815 per 1000	RR 1.15 (1.01 -1.30)	1 [107]	⊕⊖⊖ Very Low (serious imprecision;
	Difference: 106 more per 1000 (95% CI: 7 more to 213 more)			(438 patients)	serious indirectness)

95% CI: 95% Confidence interval; RR: Risk ratio

References: [12, 106, 107].

Evidence adopted: Australian guidelines for the https://app.magicapp.org/#/guideline/5446/section/78677